

THE UNITED STATES OF AMERICA

TO ALL TO WHOM THESE PRESENTS SHALL COME:

UNITED STATES DEPARTMENT OF COMMERCE
United States Patent and Trademark Office

December 05, 2001

THIS IS TO CERTIFY THAT ANNEXED HERETO IS A TRUE COPY OF
THE BELOW IDENTIFIED INTERNATIONAL APPLICATION AS
ORIGINALLY FILED AND ANY CORRECTIONS THERETO FROM THE
RECORDS OF THE UNITED STATES PATENT AND TRADEMARK
OFFICE ACTING AS A RECEIVING OFFICE UNDER THE PATENT
COOPERATION TREATY.

APPLICATION NUMBER: *PCT/US99/08516*
FILING DATE: *April 16, 1999*

By Authority of the
COMMISSIONER OF PATENTS AND TRADEMARKS



M. K. HAWKINS
Certifying Officer

HOME COPY

PCT

REQUEST

The undersigned requests that the present international application be processed according to the Patent Cooperation Treaty.

For receiving Office use only

PCT/US 99/08516

International Application No.

International Filing Date

16 APR 1999

(16.04.99)

PCT INTERNATIONAL
APPLICATION RO/US

Name of receiving Office and "PCT International Application"

Applicant's or agent's file reference

SEM4492P0142PCT^A

(if desired) (12 characters maximum)

Box No. I TITLE OF INVENTION

PROCESS AND APPARATUS FOR TREATING A WORKPIECE SUCH AS A SEMICONDUCTOR WAFER

Box No. II APPLICANT

Name and address: (Family name followed by given name; for a legal entity, full official designation)
The address must include postal code and name of country. The country of the address indicated in this
Box is the applicant's State (i.e. country) of residence if no State of residence is indicated below.)

SEMITOOL, INC.
655 Reserve Drive
Kalispell, Montana 59901
US

☐ This person is also inventor.

Telephone No.
406-751-6366

Facsimile No.
406-755-3226

Teleprinter No.

State (i.e. country) of nationality:
US

State (i.e. country) of residence:
US

This person is applicant for the purposes of: ☐ all designated States ☒ all designated States except the United States of America ☐ the United States of America only ☐ the States indicated in the Supplemental Box

Box No. III FURTHER APPLICANT(S) AND/OR (FURTHER) INVENTOR(S)

Name and address: (Family name followed by given name; for a legal entity, full official designation)
The address must include postal code and name of country. The country of the address indicated in this
Box is the applicant's State (i.e. country) of residence if no State of residence is indicated below.)

ERIC J. BERGMAN
2355 Whitefish Stage
Kalispell, Montana 59901
US

This person is:

☐ applicant only☒ applicant and inventor☐ inventor only (If this check-box is marked, do not fill in below.)

State (i.e. country) of nationality:
US

State (i.e. country) of residence:
US

This person is applicant for the purposes of: ☐ all designated States ☐ all designated States except the United States of America ☒ the United States of America only ☐ the States indicated in the Supplemental Box

☒ Further applicants and/or (further) inventors are indicated on a continuation sheet.

Box No. IV AGENT OR COMMON REPRESENTATIVE; OR ADDRESS FOR CORRESPONDENCE

The person identified below is hereby/has been appointed to act on behalf of the applicant(s) before the competent International Authorities as:

☒ agent ☐ common representative

Name and address: (Family name followed by given name; for a legal entity, full official designation. The address must include postal code and name of country.)

RANDALL T. ERICKSON
Rockey, Milnamow & Katz, Ltd.
Two Prudential Plaza, 47th Floor
180 North Stetson
Chicago, Illinois 60601
US

Telephone No.
312-616-5400

Facsimile No.
312-616-5460

Teleprinter No.

☐ Mark this check-box where no agent or common representative is/has been appointed and the space above is used instead to indicate a special address to which correspondence should be sent.

Continuation of Box No. III FURTHER APPLICANTS AND/OR (FURTHER) INVENTORS

If none of the following sub-boxes is used, this sheet is not to be included in the request.

Name and address: (Family name followed by given name; for a legal entity, full official designation)
The address must include postal code and name of country. The country of the address indicated in this
Box is the applicant's State (i.e. country) of residence if no State of residence is indicated below.)

MIGNON P. HESS
3925 South East 14th
Portland, Oregon 97202
US

This person is:

- ☐ applicant only
☒ applicant and inventor
☐ inventor only (If this check-box is marked, do not fill in below.)

State (i.e. country) of nationality:
US

State (i.e. country) of residence:
US

This person is applicant for the purposes of: ☐ all designated States ☐ all designated States except the United States of America ☒ the United States of America only ☐ the States indicated in the Supplemental Box

Name and address: (Family name followed by given name; for a legal entity, full official designation)
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Box is the applicant's State (i.e. country) of residence if no State of residence is indicated below.)

This person is:

- ☐ applicant only
☐ applicant and inventor
☐ inventor only (If this check-box is marked, do not fill in below.)

State (i.e. country) of nationality:

State (i.e. country) of residence:

This person is applicant for the purposes of: ☐ all designated States ☐ all designated States except the United States of America ☐ the United States of America only ☐ the States indicated in the Supplemental Box

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This person is:

- ☐ applicant only
☐ applicant and inventor
☐ inventor only (If this check-box is marked, do not fill in below.)

State (i.e. country) of nationality:

State (i.e. country) of residence:

This person is applicant for the purposes of: ☐ all designated States ☐ all designated States except the United States of America ☐ the United States of America only ☐ the States indicated in the Supplemental Box

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Box is the applicant's State (i.e. country) of residence if no State of residence is indicated below.)

This person is:

- ☐ applicant only
☐ applicant and inventor
☐ inventor only (If this check-box is marked, do not fill in below.)

State (i.e. country) of nationality:

State (i.e. country) of residence:

This person is applicant for the purposes of: ☐ all designated States ☐ all designated States except the United States of America ☐ the United States of America only ☐ the States indicated in the Supplemental Box

☐ Further applicants and/or (further) inventors are indicated on another continuation sheet.

Box No.V DESIGNATION OF STATES

The following designations are hereby made under Rule 4.9(a) (mark the applicable check-boxes; at least one must be marked):

Regional Patent

- ☐ **AP ARIPO Patent:** GH Ghana, KE Kenya, LS Lesotho, MW Malawi, SD Sudan, SZ Swaziland, UG Uganda, ZW Zimbabwe, and any other State which is a Contracting State of the Harare Protocol and of the PCT
- ☐ **EA Eurasian Patent:** AM Armenia, AZ Azerbaijan, BY Belarus, KG Kyrgyzstan, KZ Kazakstan, MD Republic of Moldova, RU Russian Federation, TJ Tajikistan, TM Turkmenistan, and any other State which is a Contracting State of the Eurasian Patent Convention and of the PCT
- ☒ **EP European Patent:** AT Austria, BE Belgium, CH and LI Switzerland and Liechtenstein, DE Germany, DK Denmark, ES Spain, FI Finland, FR France, GB United Kingdom, GR Greece, IE Ireland, IT Italy, LU Luxembourg, MC Monaco, NL Netherlands, PT Portugal, SE Sweden, and any other State which is a Contracting State of the European Patent Convention and of the PCT
- ☐ **OA OAPI Patent:** BF Burkina Faso, BJ Benin, CF Central African Republic, CG Congo, CI Côte d'Ivoire, CM Cameroon, GA Gabon, GN Guinea, ML Mali, MR Mauritania, NE Niger, SN Senegal, TD Chad, TG Togo, and any other State which is a member State of OAPI and a Contracting State of the PCT (if other kind of protection or treatment desired, specify on dotted line)

National Patent (if other kind of protection or treatment desired, specify on dotted line):

- | | |
|-------------------------------------------------------------------|-----------------------------------------------------------------------|
| <input type="checkbox"/> AL Albania | <input type="checkbox"/> LV Latvia |
| <input type="checkbox"/> AM Armenia | <input type="checkbox"/> MD Republic of Moldova |
| <input type="checkbox"/> AT Austria | <input type="checkbox"/> MG Madagascar |
| <input type="checkbox"/> AU Australia | <input type="checkbox"/> MK The former Yugoslav Republic of Macedonia |
| <input type="checkbox"/> AZ Azerbaijan | <input type="checkbox"/> MN Mongolia |
| <input type="checkbox"/> BA Bosnia and Herzegovina | <input type="checkbox"/> MW Malawi |
| <input type="checkbox"/> BB Barbados | <input type="checkbox"/> MX Mexico |
| <input type="checkbox"/> BG Bulgaria | <input type="checkbox"/> NO Norway |
| <input type="checkbox"/> BR Brazil | <input type="checkbox"/> NZ New Zealand |
| <input type="checkbox"/> BY Belarus | <input type="checkbox"/> PL Poland |
| <input type="checkbox"/> CA Canada | <input type="checkbox"/> PT Portugal |
| <input type="checkbox"/> CH and LI Switzerland and Liechtenstein | <input type="checkbox"/> RO Romania |
| <input checked="" type="checkbox"/> CN China | <input type="checkbox"/> RU Russian Federation |
| <input type="checkbox"/> CU Cuba | <input type="checkbox"/> SD Sudan |
| <input type="checkbox"/> CZ Czech Republic | <input type="checkbox"/> SE Sweden |
| <input type="checkbox"/> DE Germany | <input checked="" type="checkbox"/> SG Singapore |
| <input type="checkbox"/> DK Denmark | <input type="checkbox"/> SI Slovenia |
| <input type="checkbox"/> EE Estonia | <input type="checkbox"/> SK Slovakia |
| <input type="checkbox"/> ES Spain | <input type="checkbox"/> SL Sierra Leone |
| <input type="checkbox"/> FI Finland | <input type="checkbox"/> TJ Tajikistan |
| <input type="checkbox"/> GB United Kingdom | <input type="checkbox"/> TM Turkmenistan |
| <input type="checkbox"/> GE Georgia | <input type="checkbox"/> TR Turkey |
| <input type="checkbox"/> GH Ghana | <input type="checkbox"/> TT Trinidad and Tobago |
| <input type="checkbox"/> HU Hungary | <input type="checkbox"/> UA Ukraine |
| <input type="checkbox"/> IL Israel | <input type="checkbox"/> UG Uganda |
| <input type="checkbox"/> IS Iceland | <input checked="" type="checkbox"/> US United States of America |
| <input checked="" type="checkbox"/> JP Japan | <input type="checkbox"/> UZ Uzbekistan |
| <input type="checkbox"/> KE Kenya | <input type="checkbox"/> VN Viet Nam |
| <input type="checkbox"/> KG Kyrgyzstan | <input type="checkbox"/> YU Yugoslavia |
| <input type="checkbox"/> KP Democratic People's Republic of Korea | <input type="checkbox"/> ZW Zimbabwe |
| <input checked="" type="checkbox"/> KR Republic of Korea | |
| <input type="checkbox"/> KZ Kazakstan | |
| <input type="checkbox"/> LC Saint Lucia | |
| <input type="checkbox"/> LK Sri Lanka | |
| <input type="checkbox"/> LR Liberia | |
| <input type="checkbox"/> LS Lesotho | |
| <input type="checkbox"/> LT Lithuania | |
| <input type="checkbox"/> LU Luxembourg | |

Check-boxes reserved for designating States (for the purposes of a national patent) which have become party to the PCT after issuance of this sheet:

In addition to the designations made above, the applicant also makes under Rule 4.9(b) all designations which would be permitted under the PCT except the designation(s) of _____.

applicant declares that those additional designations are subject to confirmation and that any designation which is not confirmed before the expiration of 15 months from the priority date is to be regarded as withdrawn by the applicant at the expiration of that time limit. (Confirmation of a designation consists of the filing of a notice specifying that designation and the payment of the designation and confirmation fees. Confirmation must reach the receiving Office within the 15-month time limit.)

Supplemental Box *If the Supplemental Box is not used, this sheet need not be included in the request.*

Use this box in the following cases:

1. *If, in any of the Boxes, the space is insufficient to furnish all the information:*

in such case, write "Continuation of Box No. ..." [indicate the number of the Box] and furnish the information in the same manner as required according to the captions of the Box in which the space was insufficient;

in particular:

 - (i) *if more than two persons are involved as applicants and/or inventors and no "continuation sheet" is available:*

in such case, write "Continuation of Box No. III" and indicate for each additional person the same type of information as required in Box No. III;
 - (ii) *if, in Box No. II or in any of the sub-boxes of Box No. III, the indication "the States indicated in the Supplemental Box" is checked:*

in such case, write "Continuation of Box No. II" or "Continuation of Box No. III" or "Continuation of Boxes No. II and No. III" (as the case may be), indicate the name of the applicant(s) involved and, next to (each) such name, the State(s) (and/or, where applicable, ARIPO, Eurasian, European or OAPI patent) for the purposes of which the named person is applicant;
 - (iii) *if, in Box No. II or in any of the sub-boxes of Box No. III, the inventor or the inventor/applicant is not inventor for the purposes of all designated States or for the purposes of the United States of America:*

in such case write "Continuation of Box No. II" or "Continuation of Box No. III" or "Continuation of Boxes No. II and No. III" (as the case may be), indicate the name of the inventor(s) and, next to (each) such name, the State(s) (and/or, where applicable, ARIPO, Eurasian, European or OAPI patent) for the purposes of which the named person is inventor;
 - (iv) *if, in addition to the agent(s) indicated in Box No. IV, there are further agents:*

in such case, write "Continuation of Box No. IV" and indicate for each further agent the same type of information as required in Box No. IV;
 - (v) *if, in Box No. V, the name of any State (or OAPI) is accompanied by the indication "patent of addition," or "certificate of addition," or if, in Box No. V, the name of the United States of America is accompanied by an indication "Continuation" or "Continuation-in-part":*

in such case, write "Continuation of Box No. V" and the name of each State involved (or OAPI), and after the name of each such State (or OAPI), the number of the parent title or parent application and the date of grant of the parent title or filing of the parent application;
 - (vi) *if there are more than three earlier applications whose priority is claimed:*

in such case, write "Continuation of Box No. VI" and indicate for each additional earlier application the same type of information as required in Box No. VI.
2. *If the applicant claims, in respect of any designated Office, the benefits of provisions of the national law concerning non-prejudicial disclosures or exceptions to lack of novelty:*

in such case, write "Statement Concerning Non-Prejudicial Disclosures or Exceptions to Lack of Novelty" and furnish that statement below.

Box No. VI PRIORITY CLAIM

Further priority claims are indicated in the Supplemental Box ☐

The priority of the following earlier application(s) is hereby claimed:

Country (in which, or for which, the application was filed)	Filing Date (day/month/year)	Application No.	Office of filing (only for regional or international application)
item (1) US	16/April/1998 (16-04-98)	09/061,318	
item (2) US	03 September 1998 (03-09-98)	60/099,067	
item (3) US	19 March 1999 (19-03-99)	60/125,309	

Mark the following check-box if the certified copy of the earlier application is to be issued by the Office which for the purposes of the present international application is the receiving Office (a fee may be required):



The receiving Office is hereby requested to prepare and transmit to the International Bureau a certified copy of the earlier application(s) identified above as item(s):

1, 2, 3

Box No. VII INTERNATIONAL SEARCHING AUTHORITY

Choice of International Searching Authority (ISA) (If two or more International Searching Authorities are competent to carry out the international search, indicate the Authority chosen; the two-letter code may be used): ISA/US

Earlier search Fill in where a search (international, international-type or other) by the International Searching Authority has already been out or requested and the Authority is now requested to base the international search, to the extent possible, on the results of that earlier search. such search or request either by reference to the relevant application (or the translation thereof) or by reference to the search request:
Country (or regional Office): Date (day/month/year): Number:

Box No. VIII CHECK LIST

This international application contains the following number of sheets:

1. request : 5 sheets
 2. description : 28 sheets
 3. claims : 17 sheets
 4. abstract : 1 sheets
 5. drawings : 7 sheets
Total : 58 sheets

This international application is accompanied by the item(s) marked below:

1. ☐ separate signed power of attorney
 2. ☐ copy of general power of attorney
 3. ☐ statement explaining lack of signature
 4. ☐ priority document(s) identified in Box No. VI as item(s):
 5. ☒ fee calculation sheet
 6. ☐ separate indications concerning deposited microorganisms
 7. ☐ nucleotide and/or amino acid sequence listing (diskette)
 8. ☐ other (specify):

Figure No. _____ of the drawings (if any) should accompany the abstract when it is published.

Box No. IX SIGNATURE OF APPLICANT OR AGENT

Next to each signature, indicate the name of the person signing and the capacity in which the person signs (if such capacity is not obvious from reading the request).


 RANDALL T. ERICKSON, REG. NO. 33,872
 AGENT

4/16/99

1. Date of actual receipt of the purported international application: 418 Rec'd PCT/PTO 16 APR 1999		2. Drawings <input type="checkbox"/> received: <input type="checkbox"/> not received:
3. Corrected date of actual receipt due to later but timely received papers or drawings completing the purported international application:		
4. Date of timely receipt of the required corrections under PCT Article 11(2):		
5. International Searching Authority specified by the applicant: ISA/US	6. <input type="checkbox"/> Transmittal of search copy delayed until search fee is paid	

Date of receipt of the record copy by the International Bureau:

For International Bureau use only

PCT

POWER OF ATTORNEY

(for an international application filed under the Patent Cooperation Treaty)

(PCT Rule 90.4)

The undersigned applicant(s) (Names should be indicated as they appear in the request):

SEMITOOL, INC.
655 Reserve Drive
Kalispell, Montana 59901
US

hereby appoints (appoint) the following person as:



agent



common representative

Name and address

(Family name followed by given name; for a legal entity, full official designation. The address must include postal code and name of country.)

Lawrence J. Chapa	Reg. No. 39,135	Kathleen A. Lyons	Reg. No. 31,852	Elaine M. Ramesh	Reg. No. 43,032
Randall T. Erickson	Reg. No. 33,872	John P. Milnamow	Reg. No. 20,635	Keith V. Rockey	Reg. No. 24,713
Stephen D. Geimer	Reg. No. 28,846	Lisa V. Mueller	Reg. No. 38,978	Thomas I. Ross	Reg. No. 29,275
Allen J. Hoover	Reg. No. 24,103	Paul M. Odell	Reg. No. 28,332	Joel E. Siegel	Reg. No. 25,440
Martin L. Katz	Reg. No. 25,011	Robert B. Polit	Reg. No. 33,993	Paul M. Vargo	Reg. No. 29,116
Harry M. Cross, Jr.	Reg. No. 22,229				

to represent the undersigned before



all the competent International Authorities



the International Searching Authority only



the International Preliminary Examining Authority only

in connection with the international application identified below:

Title of the invention: PROCESS AND APPARATUS FOR TREATING A WORKPIECE SUCH AS A
SEMICONDUCTOR WAFER

Applicant's or agent's file reference: SEM4492P0142PC

International application number (if already available):

filed with the following Office United States as receiving Office
and to make or receive payments on behalf of the undersigned.

Signature of the applicant(s) (where there are several applicants, each of them must sign; next to each signature, indicate the name of the person signing and the capacity in which the person signs, if such capacity is not obvious from reading the request or this power):


OFFICER ON BEHALF OF SEMITOOl, INC.
W. A. Freeman

Sr. VP-Finance, CFO
TITLE

Date: 05 May 1999

PCT

POWER OF ATTORNEY

(for an international application filed under the Patent Cooperation Treaty)

(PCT Rule 90.4)

The undersigned applicant(s) (Names should be indicated as they appear in the request):

ERIC J. BERGMAN
2355 Whitefish Stage
Kalispell, Montana 59901
US

hereby appoints (appoint) the following person as:



agent



common representative

Name and address

(Family name followed by given name; for a legal entity, full official designation. The address must include postal code and name of country.)

Lawrence J. Chapa	Reg. No. 39,135	Kathleen A. Lyons	Reg. No. 31,852	Elaine M. Ramesh	Reg. No. 43,032
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Stephen D. Geimer	Reg. No. 28,846	Lisa V. Mueller	Reg. No. 38,978	Thomas I. Ross	Reg. No. 29,275
Allen J. Hoover	Reg. No. 24,103	Paul M. Odell	Reg. No. 28,332	Joel E. Siegel	Reg. No. 25,440
Martin L. Katz	Reg. No. 25,011	Robert B. Polit	Reg. No. 33,993	Paul M. Vargo	Reg. No. 29,116
Harry M. Cross, Jr. Reg. No. 22,229					

to represent the undersigned before



all the competent International Authorities



the International Searching Authority only



the International Preliminary Examining Authority only

in connection with the international application identified below:

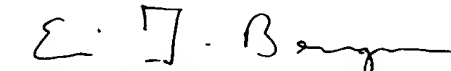
Title of the invention: PROCESS AND APPARATUS FOR TREATING A WORKPIECE SUCH AS A
SEMICONDUCTOR WAFER

Applicant's or agent's file reference: SEM4492P0142PC

International application number (if already available):

filed with the following Office United States as receiving Office
and to make or receive payments on behalf of the undersigned.

Signature of the applicant(s) (where there are several applicants, each of them must sign; next to each signature, indicate the name of the person signing and the capacity in which the person signs, if such capacity is not obvious from reading the request or this power):


ERIC J. BERGMAN

Date: 5/5/99

PCT

POWER OF ATTORNEY

(for an international application filed under the Patent Cooperation Treaty)

(PCT Rule 90.4)

The undersigned applicant(s) (Names should be indicated as they appear in the request):

MIGNON P. HESS
3925 South East 14th
Portland, Oregon 97202
US

hereby appoints (appoint) the following person as:



agent



common representative

Name and address

(Family name followed by given name; for a legal entity, full official designation. The address must include postal code and name of country.)

Lawrence J. Chapa	Reg. No. 39,135	Kathleen A. Lyons	Reg. No. 31,852	Elaine M. Ramesh	Reg. No. 43,032
Randall T. Erickson	Reg. No. 33,872	John P. Milnamow	Reg. No. 20,635	Keith V. Rockey	Reg. No. 24,713
Stephen D. Geimer	Reg. No. 28,846	Lisa V. Mueller	Reg. No. 38,978	Thomas I. Ross	Reg. No. 29,275
Allen J. Hoover	Reg. No. 24,103	Paul M. Odell	Reg. No. 28,332	Joel E. Siegel	Reg. No. 25,440
Martin L. Katz	Reg. No. 25,011	Robert B. Polit	Reg. No. 33,993	Paul M. Vargo	Reg. No. 29,116

to represent the undersigned before



all the competent International Authorities



the International Searching Authority only



the International Preliminary Examining Authority only

in connection with the international application identified below:

Title of the invention: PROCESS AND APPARATUS FOR TREATING A WORKPIECE SUCH AS A
SEMICONDUCTOR WAFER

Applicant's or agent's file reference: SEM4492P0142PC

International application number (if already available):

filed with the following Office United States as receiving Office
and to make or receive payments on behalf of the undersigned.

Signature of the applicant(s) (where there are several applicants, each of them must sign; next to each signature, indicate the name of the person signing and the capacity in which the person signs, if such capacity is not obvious from reading the request or this power):


MIGNON P. HESS

Date:

6/15/99

This sheet is not part of and does not count as a sheet of the international application.

PCT
FEE CALCULATION SHEET
Annex to the Request

For receiving Office use only
PCT/US 99/08516
International application No.

Applicant's or agent's
file reference

SEM4492P0142PCT

Date stamp of the receiving Office

(16.04.99)
16 APR 1999

Applicant
SEMITOOL, INC.

CALCULATION OF PRESCRIBED FEES

1. TRANSMITTAL FEE

240.00 **T**

\$240.

2. SEARCH FEE

450.00 **S**

\$450.

International search to be carried out by **US**

(If two or more International Searching Authorities are competent in relation to the international application, indicate the name of the Authority which is chosen to carry out the international search.)

3. INTERNATIONAL FEE

Basic Fee

The international application contains **58** sheets.
first 30 sheets

455.00 **b₁**

\$455.

28 x **\$10.00** =
remaining sheets additional amount

280.00 **b₂**

\$280.

Add amounts entered at **b₁** and **b₂** and enter total at **B**

735.00 **B**

\$735.

Designation Fees

The international application contains **6** designations.
6 x **105.00** =

630.00 **D**

\$630.

number of designation fees payable (maximum 11) amount of designation fee

Add amounts entered at **B** and **D** and enter total at **I**

1,365.00 **I**

\$1365.

(Applicants from certain States are entitled to a reduction of 75% of the international fee. Where the applicant is (or all applicants are) so entitled, the total to be entered at **I** is 25% of the sum of the amounts entered at **B** and

4. FEE FOR PRIORITY DOCUMENT

45.00 **P**

\$45.

5. TOTAL FEES PAYABLE

Add amounts entered at **T**, **S**, **I** and **P**, and enter total in the TOTAL box

2,100.00

\$2100.

TOTAL

☐ The designation fees are not paid at this time.

MODE OF PAYMENT

☐ authorization to charge
deposit account (see below)

☐ bank draft

☐ coupons

☒ cheque

☐ cash

☐ other (specify):

☐ postal money order

☐ revenue stamps

DEPOSIT ACCOUNT AUTHORIZATION (this mode of payment may not be available at all receiving Offices)

The RO/ **US** ☐ is hereby authorized to charge the total fees indicated above to my deposit account.

☒ is hereby authorized to charge any deficiency or credit any overpayment in the total fees indicated above to my deposit account.

☐ is hereby authorized to charge the fee for preparation and transmittal of the priority document to the International Bureau of WIPO to my deposit account.

04-1644

April 16, 1999

Deposit Account Number

Date (day/month/year)

Signature

- 1 -

TITLE OF THE INVENTION

PROCESS AND APPARATUS FOR TREATING A WORKPIECE
SUCH AS A SEMICONDUCTOR WAFER

5 CROSS-REFERENCE TO RELATED APPLICATIONS

The present application is a continuation-in-part of U.S.S.N. 09/061,318, filed April 16, 1998, titled "Apparatus and Method for Delivering a Treatment Liquid And Ozone To Treat The Surface Of A Workpiece" (Attorney Docket No. SEM4492P0140US), a continuation-in-part of U.S.S.N. 60/099,067, filed
10 September 3, 1998, titled "Process For Cleaning A Workpiece Such As A Semiconductor Wafer" (Attorney Docket No. SEM4492P0141US), and a continuation-in-part of U.S.S.N. __, filed __, titled "Process for Cleaning a Workpiece Such As a Semiconductor Wafer" (Attorney Docket No. SEM4492P0142US) all of which are hereby incorporated by reference.

15

STATEMENT REGARDING FEDERALLY SPONSORED
RESEARCH OR DEVELOPMENT

Not Applicable

20

BACKGROUND OF THE INVENTION

The cleaning of semiconductor wafers is often a critical step in the fabrication processes used to manufacture integrated circuits or the like. The geometries dealt with are often on the order of fractions of a micron, while the film
25 thicknesses may be on the order of 20 Angstroms. This renders the devices highly susceptible to performance degradation due to organic, particulates or

- 2 -

metallic/ionic contamination. Even silicon dioxide, which is used in the fabrication structure, can be considered a contaminant if the quality or thickness of the oxide does not meet design parameters.

Although wafer cleaning has a long history, the era of "modern" cleaning techniques is generally considered to have begun in the early 1970s when RCA developed a cleaning sequence to address the various types of contamination. Although others developed the same or similar processes in the same time frame, the general cleaning sequence in its final form is basically the same.

The first step of the RCA cleaning sequence involves removal of organic contamination using sulfuric acid and hydrogen peroxide mixtures. Ratios are typically in the range of 2:1 to 20:1, with temperatures in the range of 90 - 140 degrees Celsius. This mixture is commonly called "piranha." A recent enhancement to the removal of organic contamination replaces the hydrogen peroxide with ozone that is bubbled or injected into the sulfuric acid line.

The second step of the process involves removal of oxide films with water and HF (49%) in ratios of 200:1 to 10:1, usually at ambient temperatures. This processing typically leaves regions of the wafer in a hydrophobic condition.

The next step of the process involves the removal of particles and the re-oxidation of hydrophobic silicon surfaces using a mixture of water, hydrogen peroxide, and ammonium hydroxide, usually at a temperature of about 60 - 70 degrees Celsius. Historically, ratios of these components have been on the order of 5:1:1. In recent years, that ratio has more commonly become 5:1:0.25, or even more dilute. This mixture is commonly called "SC1" (standard clean 1) or RCA1. Alternatively, it is also known as HUANG1. Although this portion of the process

does an outstanding job of removing particles by simultaneously growing and etching away a silicon dioxide film on the surface of a bare silicon wafer (in conjunction with creating a zeta potential which favors particle removal), it has the drawback of causing metals, such as iron and aluminum, in solution to deposit on
5 the silicon surface.

In the last portion of the process, metals are removed with a mixture of water, hydrogen peroxide, and hydrochloric acid. The removal is usually accomplished at around 60 -70 degrees Celsius. Historically, ratios have been on the order of 5:1:1, but recent developments have shown that more dilute
10 chemistries are also effective, including dilute mixtures of water and HCl. This mixture is commonly referred to as "SC2" (standard clean 2), RCA2, or HUANG2.

The foregoing steps are often run in sequence, constituting what is called a "pre-diffusion clean." Such a pre-diffusion clean insures that wafers are in a highly clean state prior to thermal operations which might incorporate impurities
15 into the device layer or cause them to diffuse in such a manner as to render the device useless. Although this four-step cleaning process is considered to be the standard cleaning process in the semiconductor industry, there are many variations of the process that use the same sub-components. For example, the piranha solution may be dropped from the process, resulting in a processing sequence of:
20 HF ->SC1 ->SC2. In recent years, thin oxides have been cause for concern in device performance, so "hydrochloric acid last" chemistries have been developed. In such instances, one or more of the above-noted cleaning steps are employed with the final clean including hydrochloric acid in order to remove the silicon backside from the wafer surface.

The manner in which a specific chemistry is applied to the wafers can be as important as the actual chemistry employed. For example, HF immersion processes on bare silicon wafers can be configured to be particle neutral. HF spraying on bare silicon wafers typically shows particle additions of a few hundred
5 or more for particles at 0.2 microns nominal diameter.

Although the four-chemistry clean process described above has been effective for a number of years, it nevertheless has certain deficiencies. Such deficiencies include the high cost of chemicals, the lengthy process time required to get wafers through the various cleaning steps, high consumption of water due to
10 the need for extensive rinsing between chemical steps, and high disposal costs. The result has been an effort to devise alternative cleaning processes that yield results as good as or better than the existing four-chemistry clean process, but which are more economically attractive.

Various chemical processes have been developed in an attempt to replace
15 the existing four-chemistry cleaning process. However, such cleaning processes have failed to fully address all of the major cleaning concerns of the semiconductor processing industry. More particularly, they have failed to fully address the problem of minimizing contamination from one or more of the following contaminants: organics, particles, metals/ions, and silicon dioxide.

BRIEF SUMMARY OF THE INVENTION

A novel chemistry and application technique used in a chemical process sequence to address the contamination concerns of the microelectronic fabrication industry is set forth. These concerns are namely: minimizing contamination for organics, particles, metal/ions, and silicon dioxide. Generally stated, cleaning of workpieces, such as semiconductor wafers, is accomplished by delivery of a chemical stream to the workpiece surface. Ozone is delivered either into the liquid process stream or into the process environment. The chemical stream, which may be in the form of a liquid or vapor, is applied to the wafer in a system which enables control of the boundary layer which forms on the workpiece surface. The chemical stream may include components such as ammonium hydroxide for simultaneous particle and organic removal, another chemical to raise the pH of the solution, or other chemical additives designed to accomplish one or more specific cleaning tasks. The particular application technique is configured to develop and control a boundary layer of the liquid on the wafer surface.

In accordance with one embodiment of the process, the wafers that are to be cleaned are placed in a standard Teflon wafer cassette, or in a process chamber utilizing a "carrierless" rotor design. The latter design is common in wet processing tools, such as Spray Acid Tools and Spray Solvent Tools, available from Semitool, Inc., of Kalispell, Montana.

The cassette or wafers are placed in an enclosed environment, such as a processing chamber found in one of the foregoing tools available from Semitool, Inc.. During processing, the wafers and/or cassette are rotated in the chamber.

A processing solution is sprayed from a temperature controlled, recirculating chemical tank onto the wafer surface. This heats the surfaces of the wafers as well as the environment. If the spray is shut off, a thin liquid film remains on the wafer surfaces. However, it will generally be the case that the liquid spray is continued for the duration of the chemical process step. In the case where the wafer surface is hydrophobic, a surfactant may be added to the liquid chemical to create a thin film of liquid on the surfaces. The boundary layer of the processing solution at the wafer surface is controlled through the use of the rotation rate, the flow rate of the processing solution, and/or the injection technique (nozzle design) used to deliver the liquid (or steam) stream to the surfaces of the wafers.

Ozone is concurrently injected into the enclosed chamber during the liquid spray, either through the same manifold as the liquid delivery or through a separate manifold. Ozone injection may continue after the spray has shut off. If the wafer surfaces begin to dry (as in the case of a non-continuous spray), a brief spray is activated to replenish the liquid. This insures that the exposed wafer surfaces remain wetted at all times and that the elevated temperature at the wafer surfaces is also maintained. Generally, a continuous spray is preferred.

While ozone has a limited solubility in the hot liquid solution, it is still able to diffuse through the solution and react with the surface of the wafer (whether it is silicon, photoresist, etc.) at the liquid/solid interface. Thus diffusion, rather than dissolution, is the primary mechanism used to deliver ozone to the surfaces of the wafers. It is believed that water may help to hydrolyze carbon-carbon bonds or accelerate the oxidation of silicon surfaces by hydrolyzing silicon-hydrogen or silicon-hydroxyl bonds. The elevated temperature promotes the reaction kinetics

and the high concentration of ozone in the gas phase promotes diffusion of the ozone through the liquid film even though the increased temperature of the liquid film does not result in a solution having a high concentration of ozone dissolved in it.

5 The flow of ozone can be delivered to the process chamber through a vapor generator or the like. Such a generator is filled with water, which is temperature controlled. Thus the ozone gas stream is enriched with water vapor which maintains the boundary layer on each wafer surface at a minimal thickness so that the layer does not inhibit diffusion. At the same time, such delivery assists in
10 preventing the wafers from drying completely during the process

 In a normal spray processing operations, wafer rotational speeds are in the range of 10 -100 rpm. Such low speeds tend to allow a thick boundary layer of liquid to build up on the surfaces of the wafers to create a diffusion barrier, which, in turn, inhibits the reaction rate. It has been found, however, that a continuous
15 spray of liquid, such as the de-ionized water that is heated to maintain the surface temperature of the wafers, combined with high rotational speeds (> 300 rpm), generates a very thin boundary layer that minimizes the diffusion layer thickness thereby leading to an enhanced stripping rate. It has also been found that increases in the rotational rate of the wafers during processing results in a corresponding
20 increase in the strip rate. For example, an increase in the rotational rate from 300 to 800 rpm results in the strip rate increasing by a factor of 2 or more. A further increase to 1500 rpm has been seen to result in another two-fold increase. Rotation rates of up to 3000 rpm are anticipated.

To further enhance the disclosed process, the temperature of the liquid supply (water supply) is heated to generate a supply of saturated steam under pressure to the process chamber. Under such circumstances, it is possible to achieve wafer surface temperatures in excess of 100 degrees Celsius, thereby
5 further accelerating the reaction kinetics. A steam generator may be used to pressurize the process chamber to achieve the desired temperatures. For example, saturated steam at 126 degrees Celsius may be used with a corresponding increase in the pressure of the process chamber to 35 psia. The increased pressure within the processing chamber also provides for use of higher ozone concentrations, thereby
10 generating a higher diffusion gradient across the boundary layer at the surface of each wafer. Still further, the use of steam also allows for the use of lower rotation rates to achieve the requisite thin boundary layers at the surfaces of the wafers. The oxidation rate of the ozone may also be enhanced by irradiating the surfaces of the wafers with ultra-violet light.

15 The present inventors have also found that it is now possible to remove particles, metals, and organics in a concurrent manner in a single processing step. Further, it is now possible to regenerate a fresh, clean, controlled chemical oxide film in that same step. To this end, certain additives may be provided in the processing liquid to specifically target certain contaminants and/or to enhance the
20 effectiveness of the overall process. For example, ammonium hydroxide may be added to the processing liquid (e.g., deionized water) to reduce particle counts on the workpieces. In such a process, the ozone prevents pitting of the silicon surface by the ammonium hydroxide.

Other additives that enhance the cleaning capability of the overall process include HF and HCl. Such additives have the following benefits/effects: 1) removal of organic contaminants; 2) removal of oxide and regeneration of a controlled chemical oxide; 3) removal of particles; 4) removal of metals.

5 After one or more of the foregoing cleaning process steps has been completed, the wafers are prepared for subsequent cleaning steps. To this end, the wafers are subject to a rinsing with deionized water or a suitable aqueous solution. At this time, the ozone within the processing chamber may also be purged with, for example, a nitrogen flush.

10 In instances in which an additive that enhances the metal removal capabilities of the solution is not used, it may be desirable to execute a further processing step for metal removal. In one or more such cleaning steps, metal and/or silicon dioxide may be removed from the surfaces of the wafers by applying a temperature controlled mixture containing hydrofluoric acid and/or hydrochloric
15 acid, chloroacetic acid, or other halogenated chemistry. Ozone may or may not be introduced into the liquid stream or the process environment during this step.

After one or more of the foregoing steps have been completed, including any intermediate cleaning steps, the wafers are subject to a final rinsing in deionized water or an aqueous solution. After the rinse, the wafers are dried in a
20 manner that may include the use of heated nitrogen, another inert gas flow, or organic vapors. Additionally, the wafers may be rotated during the drying process.

The disclosed process is applicable to a number of situations that require cleaning or selective removal of contaminants from the surface of a workpiece. For example, one or more of the disclosed process steps may be used to remove

photoresist from the surface of a semiconductor wafer. Still further, it is now possible to concurrently remove a layer of photoresist and a corresponding layer of an anti-reflective coating (ARC) in a single processing step using a single processing solution. More particularly, the present inventor has found that an aqueous solution having a high pH, such as a solution of ammonium hydroxide and/or tetra-methyl ammonium hydroxide and deionized water, may be used to form a controlled boundary layer that cooperate with ozone to remove both the photoresist and the anti-reflective coating. From the foregoing description, there are several novel aspects to the disclosed process. They include:

- 10 1) The use of a temperature controlled liquid chemical source delivered to the wafer surface to stabilize the temperature of the wafer and, depending on the liquid utilized, provide a supply of water to support hydrolysis of the carbon-carbon bonds of contaminants at the surface of each wafer.
- 15 2) The control of the thickness of the boundary layer of liquid present on the wafer surface so that it is not of sufficient thickness to significantly inhibit the diffusion of ozone to the wafer surface. As such, the ozone is allowed to diffuse through the controlled boundary layer, where it can oxidize silicon, organics, or metals at the surface, or otherwise support any desired reaction. The boundary layer may be controlled through the control of wafer rotation rate, vapor delivery, controlled liquid spray, the use of steam, the use of
20 surfactants or a combination of more than one of these techniques.
- 3) The process takes place in an enclosed processing chamber, which may or may not be used to produce a pressurized processing environment.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWINGS

Fig. 1 is a schematic block diagram of one embodiment of an apparatus for
5 treating a semiconductor workpiece in which ozone is injected into a line
containing a pressurized treatment liquid.

Fig. 2 is a schematic block diagram of one embodiment of an apparatus for
treating a semiconductor workpiece in which the semiconductor workpiece is
indirectly heated by heating a treatment liquid that is sprayed on the surface of the
10 workpiece.

Fig. 3 is a flow diagram illustrating one embodiment of a process flow for
treating a semiconductor workpiece with a treatment fluid and ozone.

Fig. 4 is a schematic block diagram of an alternative embodiment of the
system set forth in Fig. 2 wherein the ozone and treatment fluid are provided to the
15 semiconductor workpiece along different flow paths.

Fig. 5 is a schematic block diagram of an embodiment of an apparatus for
treating a semiconductor workpiece in which pressurized steam and ozone are
provided in a pressurized chamber containing a semiconductor workpiece.

Fig. 6 is a schematic block diagram of an embodiment of an apparatus for
20 treating a semiconductor workpiece in which an ultra-violet lamp is used to
enhance the kinetic reactions at the surface of the workpiece.

Fig. 7 is a schematic block diagram of an embodiment of an apparatus for
treating a semiconductor workpiece in which liquid gas contactors are used to
enhance the kinetic reactions at the surface of the workpiece.

DETAILED DESCRIPTION OF THE INVENTION

Apparatus for supplying a mixture of a treatment liquid and ozone for treatment of a surface of a workpiece, such as a semiconductor workpiece, to execute the foregoing processes are set forth below. The preferred embodiment of the apparatus comprises a liquid supply line that is used to provide fluid communication between a reservoir containing the treatment liquid and a treatment chamber housing the semiconductor workpiece. A heater is disposed to heat the workpiece, either directly or indirectly. Preferably, the workpiece is heated by heating the treatment liquid that is supplied to the workpiece. One or more nozzles accept the treatment liquid from the liquid supply line and spray it onto the surface of the workpiece while an ozone generator provides ozone into an environment containing the workpiece.

One embodiment of an apparatus suitable for providing ozone and a treatment liquid for treatment of a semiconductor workpiece in accordance with the four or processes is illustrated in Fig. 1. The treatment system, shown generally at 10, includes a treatment chamber 15 that contains one or more workpieces 20, such as semiconductor wafer workpieces. Although the illustrated system is directed to a batch workpiece apparatus, it will be recognized that the system is readily adaptable for use in single workpiece processing as well.

The semiconductor workpieces 20 are supported within the chamber 15 by one or more supports 25 extending from, for example, a rotor assembly 30. Rotor assembly 30 seals with the housing of the treatment chamber 15 to form a sealed, closed processing environment. Further, rotor assembly 30 is provided so that the

semiconductor workpieces 20 may be spun about axis 35 during or after treatment with the ozone and treatment liquid.

One or more nozzles 40 are disposed within the treatment chamber 15 so as to direct a spray mixture of ozone and treatment liquid onto the surfaces of the semiconductor workpieces 20 that are to be treated. In the illustrated embodiment, the nozzles 40 direct a spray of treatment fluid to the underside of the semiconductor workpieces 20. However, it will be recognized that the fluid spray may be directed alternatively, or in addition, to the upper surface of the semiconductor workpieces 20.

Treatment liquid and ozone are supplied to the nozzles 40 with the assistance of a number of system components that are uniquely arranged to provide a single fluid line comprising ozone mixed with the treating liquid. To this end, a reservoir 45 defines a chamber 50 in which the liquid that is to be mixed with the ozone is stored. The chamber 50 is in fluid communication with the input of a pump mechanism 55. The pump mechanism 55 provides the liquid under pressure along a fluid flow path, shown generally at 60, for ultimate supply to the input of the nozzles 40. The preferred treatment fluid is deionized water, but it will be recognized that other treatment fluids, such as other aqueous or non-aqueous solutions, may also be employed.

A number of components are disposed along the fluid flow path 60. First, a filter 65 is disposed along the fluid flow path 60 to filter out microscopic contaminants from the treatment fluid. The treatment fluid, still under pressure, is provided at the output of the filter 65 along fluid flow line 70. It is along fluid flow line 70 that ozone is injected. The ozone is generated by ozone generator 75 and is

supplied along fluid flow line 80 under pressure to fluid flow line 70. Optionally, the treatment liquid, now injected with ozone, is supplied to the input of a mixer 90 that mixes the ozone and the treatment liquid. The mixer 90 may be static or active. From the mixer 90, the treatment liquid and ozone are provided to be input of
5 nozzles 40 which, in turn, spray the liquid on the surface of the semiconductor workpieces 20 that are to be treated and, further, introduce the ozone into the environment of the treatment chamber 15.

To further concentrate the ozone in the treatment liquid, an output of the ozone generator 75 may be supplied to a dispersion unit 95 disposed in the liquid
10 chamber 50 of the reservoir 45. The dispersion unit 95 provides a dispersed flow of ozone through the treatment liquid to thereby add ozone to the fluid stream prior to injection of a further amount of ozone along the fluid path 60.

In the embodiment of the system of Fig. 1, spent liquid in chamber 15 is provided along fluid line 105 to, for example, a valve mechanism 110. The valve
15 mechanism 110 may be operated to provide the spent liquid to either a drain output 115 or back to the liquid chamber 50 of the reservoir 45. Repeated cycling of the treatment liquid through the system and back to the reservoir 45 assists in elevating the ozone concentration in the liquid through repeated ozone injection and/or ozone dispersion.

20 A further embodiment of a system for delivering a fluid mixture for treating the surface of a semiconductor workpiece is illustrated in Fig. 2. Although the system 120 of Fig. 2 appears to be substantially similar to the system 10 of Fig. 1, there are significant differences. The system 120 of Fig. 2 is based upon the recognition by the present inventors that the heating of the surfaces of the

semiconductor workpieces 20 with a heated liquid that is supplied along with a flow of ozone that creates an ozonated atmosphere is highly effective in photoresist stripping, ash removal, and/or cleaning processes. As such, system 120 includes one or more heaters 125 that are used to heat the treatment liquid so that it is
5 supplied to the surfaces of the semiconductor workpieces at an elevated temperature that accelerates the surface reactions. It will be recognized that it is also possible to directly heat the workpieces so as to stimulate the reactions. Such heating may take place in addition to or instead of the indirect heating of the workpieces through contact with the heated treatment liquid. For example,
10 supports 25 may include heating elements that may be used to heat the workpieces 20. The chamber 15 may include a heater for elevating the temperature of the chamber environment and workpieces.

As noted above, the preferred treatment liquid is deionized water since it is believed to be required to initiate the cleaning/removal reactions at the workpiece
15 surface, probably through hydrolysis of the carbon-carbon bonds of organic molecules. The present inventors, however, recognize that significant amounts of water can form a continuous film on the semiconductor workpiece surface. This film acts as a diffusion barrier to the ozone, thereby inhibiting reaction rates. Control of the boundary layer thickness, as will be explained in further detail
20 below, is implemented by control of the rpm of the semiconductor workpiece, vapor delivery, and controlled spraying of the treatment liquid, or a combination of one or more of these techniques. By reducing the boundary layer thickness, the ozone is allowed to diffuse to the surface of the workpieces and react with the organic materials that are to be removed.

Fig. 3 illustrates one embodiment of a process that may be implemented in the system of Fig. 2 when the system 120 is used, for example, to strip photoresist from the surfaces of semiconductor workpieces. At step 200, the workpieces 20 that are to be stripped are placed in, for example, a Teflon wafer cassette. This
5 cassette is placed in a closed environment, such as in chamber 15. Chamber 15 and its corresponding components may be constructed based on a spray solvent tool platform or spray acid tool platform such as those available from Semitool, Inc., of Kalispell, Montana. Alternatively, the semiconductor workpieces 20 may be disposed in chamber 15 in a carrierless manner, consistent with the automated
10 processing platform design of the MAGNUM® brand semiconductor processing tool available from Semitool, Inc.

At step 205, heated deionized water is sprayed onto the surfaces of the semiconductor workpieces 20. The heated deionized water heats the surfaces of the semiconductor workpieces 20 as well as the enclosed environment of the
15 chamber 15. When the spray is discontinued, a thin liquid film remains on the workpiece surfaces. If the surface is hydrophobic, a surfactant may be added to the deionized water to assist in creating a thin liquid boundary layer on the workpiece surfaces. In view of the teachings set forth herein, it will be recognized that a surfactant may be used in connection with hydrophilic surfaces as well. Corrosion
20 inhibitors may also be used in conjunction with the aqueous ozone, thin boundary layer process.

The surface boundary layer of deionized water is controlled at step 210 using one or more techniques. For example, the semiconductor workpieces 20 may be rotated about axis 35 by rotor 30 to thereby generate centripetal

accelerations that thin the boundary layer. The flow rate of the deionized water may also be used to control the thickness of the surface boundary layer. Lowering of the flow rate results in decreased boundary layer thickness. Still further, the manner in which the deionized water is injected into the chamber 15 may be used to control the boundary layer thickness. Nozzles 40 may be designed to provide the deionized water as micro-droplets thereby resulting in a thin boundary layer.

At step 215, ozone is injected into the fluid flow path 60 during the water spray, or otherwise provided to the internal chamber environment of chamber 15. If the apparatus of Fig. 2 is utilized, the injection of the ozone continues after the spray has shut off. If the workpiece surface begins to dry, a brief spray is preferably activated to replenish the liquid film on the workpiece surface. This ensures that the exposed workpiece surfaces remain wetted at all times and, further, ensures that the workpiece temperature is and remains elevated at the desired reaction temperature. It has been found that a continuous spray of deionized water having a flow rate that is sufficient to maintain the workpiece surfaces at an elevated temperature, and high rotational speeds (i.e., > 300 rpm, between 300 and 800 rpm, or even as high as or greater than 1500 rpm) generates a very thin boundary layer which minimizes the ozone diffusion barrier and thereby leads to an enhanced photoresist stripping rate. As such, the control of the boundary layer thickness is used to regulate the diffusion of reactive ozone to the surface of the wafer.

While ozone has a limited solubility in the heated deionized water, the ozone is able to diffuse through the water and react with photoresist at the liquid/resist interface. It is believed that the presence of the deionized water itself

further assists in the reactions by hydrolyzing the carbon-carbon bonds of organic deposits, such as photoresist, on the surface of the wafer. The higher temperature promotes the reaction kinetics while the high concentration of ozone in the gas phase promotes diffusion of ozone through the boundary layer film even though
5 the high temperature of the boundary layer film does not actually have a high concentration of dissolved ozone.

After the semiconductor workpieces 20 have been processed through the reactions of the ozone and/or liquid with the materials to be removed, the workpieces are subject to a rinse at 220 and are dried at step 225. For example, the
10 workpieces may be sprayed with a flow of deionized water during the rinse at step 220. They may then be subject to any one or more known drying techniques thereafter at step 225.

In accordance with the foregoing process, elevated temperatures are used to accelerate the reaction rates at the wafer surface. One manner in which the surface
15 temperature of the wafer may be maximized is to maintain a constant delivery of heated processing liquid, such as water or steam, during the process. The heated processing liquid contacts and heats the wafer during processing. However, such a constant delivery may result in significant waste of the water or other processing liquid. In order to conserve water and achieve the thinnest possible boundary
20 layer, a "pulsed flow" of liquid or steam may be used. In instances in which such a "pulsed flow" fails to maintain the requisite elevated wafer surface temperatures, an alternative manner of maintaining the wafer surface temperature may be needed. One such alternative is the use of a "hot wall" reactor that maintains the wafer surface and processing environment temperatures at the desired level. To

this end, the process chamber may be heated by, for example, one or more embedded heated recirculating coils, a heating blanket, irradiation from a thermal source (e.g., and infrared lamp), etc.

In laboratory experiments, a 150 mm silicon wafer coated with 1 micron of photoresist was stripped in accordance with the teachings of the foregoing process. The processing chamber was pre-heated by spraying deionized water that was heated to 95 degrees Celsius into the processing chamber for 10 minutes. During the cleaning process, a pulsed flow of deionized water heated to 95 degrees Celsius was used. The pulsed flow included an "on time" of approximately five seconds followed by an "off time" of 10 seconds. The wafer was rotated at 800 rpm and the pulsed flow of deionized water was sprayed into the processing chamber through nine nozzles at a rate of 3 liters per minute. Ozone was injected into the processing chamber through a separate manifold at a rate of 8 liters per minute at a concentration of 12 percent. The resultant strip rate was 7234 Angstroms/min.

There are many benefits resulting from the use of the semiconductor cleaning processes described above. One of the most significant benefits is that the conventional 4-chem clean process may be reduced to a two-chemical step process while retaining the ability to remove organics, remove particulates, reduce metals and remove silicon dioxide. Process times, chemical consumption, water consumption and waste generation are all also significantly reduced. A further benefit of the foregoing process is its applicability to both FEOL and BEOL wafers and strip processes. Laboratory tests indicate that there is no attack on metals such as aluminum, titanium, tungsten, etc. A known exception is copper, which forms a copper oxide in the presence of ozone. This oxide is not a "hard" and uniform

passivation oxide, such as the oxide that forms on metals like aluminum. As a result, the oxide can be readily removed.

With reference again to Figure 3, it will be recognized that process steps 205 - 215 may be executed in a substantially concurrent manner. Additionally, it will be recognized that process steps 205 - 215 may be sequentially repeated using different processing liquids. In such instances, each of the processing liquids that are used may be specifically tailored to remove a respective set of contaminants. Preferably, however, it is desirable to use as few different processing liquids as possible. By reducing the number of different processing liquids utilized, the overall cleaning process is simplified and reducing the number of different processing liquids utilized minimizes chemical consumption.

The present inventors have developed a single processing liquid that may be utilized to remove organic contaminants, metals, and particles in a single cycle of process steps 205-215. The processing liquid is comprised of a solution of deionized water and one or more compounds, such as HF or HCl, so as to form an acidic processing liquid solution.

It has now been shown that the use of a hydrofluoric acid solution in the process steps set forth at 205-215 provides numerous advantages, including the following:

1. Removal of organic contaminants - The oxidation capability of the process has been demonstrated repeatedly on photoresist. Strip rates often exceed 4000A/minute. Considering the fact that in cleaning applications, organic contamination is generally on the molecular level, the disclosed process has ample oxidation capacity.

2. Removal of oxide and regeneration of a controlled chemical oxide -
Depending on the temperature of the solution and the concentration of HF
in solution, a specific etch rate may be defined. However, the ozone will
diffuse through the controlled boundary layer and regenerate the oxide to
prevent the wafer from becoming hydrophobic. A 500:1 H₂O:HF mixture at
65 degrees C will etch SiO₂ at a rate of about 6A/minute. The same solution
at 25 degrees C will etch SiO₂ at about 2A/minute. A typical "native" oxide
is generally self limiting at a thickness of 8 - 12A, which is generally the
targeted thickness for the oxide removal.
3. Removal of particles - Although the acidic solutions do not have the
favorable zeta potential present in the SC1 clean noted above, particle
removal in the disclosed process with an HF processing liquid has still been
shown to be significant, as it uses the same removal mechanism of etching
and regenerating the oxide surface,
4. Removal of metals - In laboratory experiments, wafers were intentionally
contaminated with iron, nickel and copper. The disclosed process with an
HF containing processing liquid showed a reduction in metals of over three
orders of magnitude. As an added enhancement, HCl can be used in place
of the HF to accomplish the metals removal, although this does not have
the same degree of oxide and particle removal capability. The combination
of HF and HCl is a further benefit, as each of these chemistries has
significant metals removal capability, but the regeneration of the oxide
surface in conjunction with the conversion of metals to metallic oxides and

the symbiotic interaction of the two acid halides creates an exceptionally favorable environment for metal removal.

- 5 5. An oxide-free (hydrophobic) surface may be generated, if desired, by using a final HF step in an immersion cell or by use of an HF vapor step after the metals removal.

Typical chemical application times are in the range of 1:00 to 5:00. Compared to a 4-chem clean process time of around 20:00, the disclosed process with an HF and/or HCl containing processing liquid becomes very attractive. Typical H₂O:HF:HCl concentration ratios are on the order of 500: 1:1 to 50: 1: 1, 10 with and without HF and/or HCl. Higher concentrations are possible, but the economic benefits are diminished. It is important to note that gaseous HF or HCl could be injected into water to create the desired cleaning chemistry as well. Due to differences in processor configurations and desired cleaning requirements, definition of specific cleaning process parameters will vary based on these 15 differences and requirements.

The process benefits include the following:

1. Reduction in the amount and types of chemicals used in the cleaning process.
2. Reduction in water consumption by the elimination of the numerous 20 intermediate rinse steps required.
3. Reduction in process time.
4. Simplification of process hardware.

The disclosed processes are counter-intuitive. Efforts have been made for a number of years to replace hydrogen peroxide with ozone in chemistries such as

SC1 and, to a lesser degree, SC2. These efforts have largely failed because they have not controlled the boundary layer and have not introduced the ozone in such a manner that diffusion through the boundary layer is the controlling mechanism instead of dissolution into the boundary layer. While the cleaning efficiency of conventional solutions is greatly enhanced by increasing temperature, it is recognized that the solubility of ozone in a given liquid solution is inversely proportional to the temperature of the solution. The solubility of ozone in water at 1 degrees Celsius is approximately 100 ppm. At 60 degrees Celsius, this solubility drops to less than 5 ppm. At elevated temperatures, the ozone concentration is thus insufficient to passivate (oxidize) a silicon wafer surface quickly enough to ensure that pitting of the silicon surface will not occur. Thus the two mechanisms are in conflict with one another when attempting to optimize process performance.

Tests have demonstrated that by applying the boundary layer control techniques explained in connection with the presently disclosed processes, it is possible to process silicon wafers using a 4:1 water:ammonium hydroxide solution at 95C and experience an increase surface roughness (RMS) of less than 2 angstroms. When this same solution is applied in an immersion system or in a conventional spray system, RMS surface roughness as measured by atomic force microscopy increases by more than 20 angstroms and the maximum surface roughness exceeds 190 angstroms. Additionally, while a conventional process will pit the surface to such a degree as to render the surface unreadable by a light-scattering particle counter, the boundary controlled technique has actually shown particle reductions of up to 50% on the wafer surface.

In the case of oxidizing and removing organic contamination, conventional aqueous ozone processes show a strip rate on photoresist (a hydrocarbon film) of around 200 - 700 angstroms per minute. In the boundary layer controlled system of the disclosed processes, the rate is accelerated to 2500 to 7000 angstroms per minute in a spray controlled boundary layer, or 8000 angstroms per minute when the boundary layer is generated and controlled using steam at 15 psi and 126 degrees C.

The disclosed processes are suitable for use in a wide range of microelectronic fabrication applications. One issue which is of concern in the manufacture of semiconductor devices is reflective notching. In order to expose a pattern on a semiconductor wafer, the wafer is coated with a photo-active compound called photoresist. The resistance film is exposed to a light pattern, thereby "exposing" the regions to which the light is conveyed. However, since topographic features may exist under the photoresist, it is possible for the light to pass through the photoresist and reflect off of a topographic feature. This results in resist exposure in an undesirable region. This phenomenon is known as "reflective notching." As device density increases, reflective notching becomes more of a problem.

A similar issue arises as a result of the reflectance normal to the incident angle of irradiation. Such reflectance can create distortions in the exposure beam through the phenomenon of standing wave formation, thereby resulting in pattern distortion in the photoresist.

In order to combat these phenomena, the use of anti-reflective coating layers has become common. The photoresist films are typically deposited either on

top of or below an anti-reflective coating layer. Since both the photoresist layer and the anti-reflective coating layer are merely "temporary" layers used in intermediate fabrication steps, they must be removed after such intermediate fabrication steps are completed.

5 It has been found that the process of Figure 3 may be used with a processing liquid comprised of water and ammonium hydroxide to remove both the photoresist and the anti-reflective coating in a single processing step (e.g., the steps illustrated at 210-215). Although this has been demonstrated at concentrations between 0.02% and 0.04% ammonium hydroxide by weight in water, other
10 concentrations are also considered to be viable.

 The process for concurrently removing photoresist and the corresponding anti-reflective layer is not necessarily restricted to processing liquids that include ammonium hydroxide. Rather, the principal goal of the additive is to elevate the pH of the solution that is sprayed onto the wafer surface. Preferably, the pH
15 should be raised so that it is between about 8.5 and 11. Although bases such as sodium hydroxide and/or potassium hydroxide may be used for such removal, they are deemed to be less desirable due to concerns over mobile ion contamination. However, chemistries such as TMAH (tetra-methyl ammonium hydroxide) are suitable and do not elicit the same a mobile ion contamination concerns. Ionized
20 water that is rich in hydroxyl radicals may also be used.

 The dilute ammonium hydroxide solution may be applied in the process in any number of manners. For example, syringe pumps, or other precision chemical applicators, can be used to enable single-use of the solution stream. In such an embodiment, it becomes possible to strip the photoresist using a deionized water

stream with ozone, and can conclude the strip with a brief period during which ammonium hydroxide is injected into the aqueous stream. This assists in minimizing chemical usage and waste generation. The application apparatus may also be capable of monitoring and controlling the pH the using the appropriate
5 sensors and actuators, for example, by use of microprocessor control.

With reference to Fig. 4, there is shown yet a further embodiment of the ozone treatment system 227. In the embodiment of Fig. 4, one or more nozzles 230 are disposed within the treatment chamber 15 to conduct ozone from ozone generator 75 directly into the reaction environment. The heated treatment fluid is
10 provided to the chamber 15 through nozzles 40 that receive the treatment fluid, such as heated deionized water, through a supply line that is separate from the ozone supply line. As such, injection of ozone in fluid path 60 is optional.

Another embodiment of an ozone treatment system is shown generally at 250 in Fig. 5. In the system 250, a steam boiler 260 that supplies saturated steam
15 under pressure to the process chamber 15 has replaced the pump mechanism. The reaction chamber 15 is preferably sealed to thereby form a pressurized atmosphere for the reactions. For example, saturated steam at 126 degrees Celsius could be generated by steam boiler 260 and supplied to reaction chamber 15 to generate a pressure of 35 psia therein during the workpiece processing. Ozone may be directly
20 injected into the chamber 15 as shown, and/or may be injected into the path 60 for concurrent supply with the steam. Using the system architecture of this embodiment, it is thus possible to achieve semiconductor workpiece surface temperatures in excess of 100 degrees Celsius, thereby further accelerating the reaction kinetics.

A still further enhancement that may be made to any one of the foregoing systems is illustrated in Fig. 6. In this embodiment, an ultra-violet lamp 300 is used to irradiate the surface of the semiconductor workpiece 20 during processing. Such irradiation further enhances the reaction kinetics. Although this irradiation technique is applicable to batch semiconductor workpiece processing, it is more easily and economically implemented in the illustrated single wafer processing environment where the workpiece is more easily completely exposed to the UV radiation.

With reference to Fig. 7, a further system 310 for implementing one or more of the foregoing processes is set forth. Of particular note in system 310 is the use of one or more liquid-gas contactors 315 that are used to promote the dissolution of ozone into the aqueous stream. Such contactors are of particular benefit when the temperature of the processing liquid is, for example, at or near ambient. Such low temperatures may be required to control corrosion that may be promoted on films such as aluminum/silicon/copper.

The contactor 315 is preferably of a parallel counter-flow design in which liquid is introduced into one end and the ozone gas is introduced into the opposite end. Such contactors are manufactured and marketed by the W. L. Gore, Corp., as well as other vendors. These contactors operate under pressure, typically from about 1 to 4 atmospheres (gauge). The undissolved gas exiting the contactor 315 may be optionally directed to the process chamber 320 to minimize gas losses. However, the ozone supply 330 for the contactor 315 may or may not be the same as the supply for direct delivery to the process chamber 320.

The presently disclosed apparatus and methods may be used to treat workpieces beyond the semiconductor workpieces described above. For example, other workpieces, such as flat panel displays, hard disk media, CD glass, etc, may also be have their surfaces treated using the foregoing apparatus and methods.

5 Although the preferred treatment liquid for the disclosed application is deionized water, other treatment liquids may also be used. For example, acidic and basic solutions may be used, depending on the particular surface to be treated and the material that is to be removed. Treatment liquids comprising sulfuric acid, hydrochloric acid, and ammonium hydroxide may be useful in various
10 applications.

Numerous modifications may be made to the foregoing system without departing from the basic teachings thereof. Although the present invention has been described in substantial detail with reference to one or more specific embodiments, those of skill in the art will recognize that changes may be made
15 thereto without departing from the scope and spirit of the invention as set forth in the appended claims.

WHAT IS CLAIMED IS:

1. A method for treating a workpiece comprising the steps of:

5 providing a heated liquid onto the surface of the workpiece that is to be
 treated, the heated liquid assisting in maintaining the workpiece at
 an elevated temperature;
 introducing an amount of ozone into an environment containing the
 workpiece;
 controlling the thickness of the heated liquid on the workpiece so as to form
10 a thin liquid boundary layer that allows for diffusion of the ozone
 therethrough for reaction at the surface of the workpiece.

2. A method for treating a workpiece as claimed in claim 1 wherein the liquid is
 deionized water.
15
3. A method for treating a workpiece as claimed in claim 2 wherein the deionized
 water is superheated.
4. A method for treating a workpiece as claimed in claim 1 wherein the liquid
20 comprises sulfuric acid.
5. A method for treating a workpiece as claimed in claim 1 wherein the liquid
 comprises hydrochloric acid.

6. A method for treating a workpiece as claimed in claim 1 wherein the liquid comprises an acid.
7. A method for treating a workpiece as claimed in claim 1 wherein the liquid
5 comprises ammonium hydroxide.
8. A method for treating a workpiece as claimed in claim 1 wherein the step of controlling comprises the step of rotating the workpiece.
- 10 9. A method for treating a workpiece as claimed in claim 1 wherein the step of controlling comprises the step of rotating the workpiece at a rotation rate equal to or greater than about 300 rpm.
10. A method for treating a workpiece as claimed in claim 1 wherein the step of
15 controlling comprises the step of adding a surfactant to the liquid.
11. A method for treating a workpiece as claimed in claim 1 wherein the step of controlling comprises the step of controlling the flow of the liquid sprayed onto the surface of the wafer that is to be treated.
- 20 12. A method for cleaning the surface of a workpiece comprising the steps of:
providing a heated liquid solution of water and a further compound selected from the group consisting of HF and HCl onto the surface of the workpiece, the heated liquid solution being suitable for concurrent

removal of organic contaminants, metals, and particles, the heated solution assisting in maintaining the workpiece at an elevated temperature;

introducing an amount of ozone into an environment containing the workpiece;

controlling the thickness of the heated liquid solution to form a thin liquid boundary layer on the surface of the workpiece to allow diffusion of the ozone therethrough for reaction at the surface of the workpiece.

10 13. A method for cleaning as claimed in claim 12 wherein the step of controlling comprises the step of rotating the workpiece at a rotation rate approximately equal to or greater than about 300 rpm.

15 14. A method for cleaning as claimed in claim 12 wherein the heated liquid is provided as a pulsed stream.

15. A method for cleaning as claimed in claim 12 wherein the liquid comprises a solution of water and HF.

20 16. A method for cleaning as claimed in claim 15 wherein the concentration ratio of water to HF is between about 50: 1 and 500: 1.

17. A method for cleaning as claimed in claim 12 wherein the liquid comprises a solution of water and HCl.

18. A method for cleaning as claimed in claim 17 wherein the concentration ratio of water to HCl is between about 50: 1 and 500: 1.

5 19. A method for cleaning as claimed in claim 12 wherein the liquid comprises a solution of water, HF, and HCl.

20. A method for cleaning as claimed in claim 19 wherein the concentration ratio of water to HF to HCl is between about 50: 1: 1 and 500: 1: 1.

10

21. A method for concurrently removing photoresist and an anti-reflective coating material from a workpiece comprising the steps of:

providing a heated liquid onto the surface of the workpiece to contact the photoresist and the anti-reflective coating material, the heated liquid comprising water and ammonium hydroxide, the heated liquid assisting in maintaining the workpiece at an elevated temperature; introducing an amount of ozone into an environment containing the workpieces;

15

controlling the thickness of the heated liquid on the surface of the workpiece to form a thin liquid boundary layer that allows diffusion of the ozone therethrough for reaction with the photoresist and antireflective coating material.

20

22. A method as claimed in claim 21 wherein the step of controlling comprises the step of rotating the workpiece at a rotation rate approximately equal to or greater than about 300 rpm.

5 23. A method for cleaning as claimed in claim 12 wherein the heated liquid is provided as a pulsed stream.

24. A method for concurrently removing an oxide from the surface of a workpiece and regenerating a controlled chemical oxide on the surface of the workpiece,
10 the method comprising the steps of:

providing a heated liquid solution of water and a further compound selected from the group consisting of HF and HCl onto the surface of the workpiece to contact the oxide, the heated liquid solution assisting in removal of existing oxide and in maintaining the workpiece at an
15 elevated temperature;

introducing an amount of ozone into an environment containing the workpiece;

controlling the thickness of the heated liquid solution to form a thin liquid boundary layer on the surface of the workpiece to allow diffusion of
20 the ozone therethrough for regeneration of a controlled oxide layer at the surface of the workpiece.

25. A method as claimed in claim 24 wherein the step of controlling comprises the step of rotating the workpiece at a rotation rate approximately equal to or greater than about 300 rpm.
- 5 26. A method as claimed in claim 24 wherein the heated liquid is provided as a pulsed stream.
27. A method as claimed in claim 24 wherein the liquid comprises a solution of water and HF.
- 10 28. A method as claimed in claim 27 wherein the concentration ratio of water to HF is between about 50: 1 and 500: 1.
29. A method as claimed in claim 24 wherein the liquid comprises a solution of
- 15 water and HCl.
30. A method as claimed in claim 29 wherein the concentration ratio of water to HCl is between about 50: 1 and 500: 1.
- 20 31. A method as claimed in claim 24 wherein the liquid comprises a solution of water, HF, and HCl.
32. A method as claimed in claim 31 wherein the concentration ratio of water to HF to HCl is between about 50: 1: 1 and 500: 1: 1.

33. A method for treating a workpiece comprising the steps of:
- providing a flow of a heated pressurized liquid;
 - 5 injecting an amount of ozone into the pressurized liquid flow;
 - spraying the mixture of ozone and heated liquid onto the workpiece.
34. A method for treating a workpiece as claimed in claim 33 wherein the liquid is
- 10 deionized water.
35. A method for treating a workpiece as claimed in claim 34 wherein the
- deionized water is heated.
36. A method for treating a workpiece as claimed in claim 34 wherein the
- 15 deionized water is superheated.
37. A method for treating a workpiece as claimed in claim 33 wherein the liquid
- comprises an acid.
- 20 38. A method for treating a workpiece as claimed in claim 33 wherein the liquid
- comprises sulfuric acid.
39. A method for treating a workpiece as claimed in claim 33 wherein the liquid
- comprises hydrochloric acid.

40. A method for treating a workpiece as claimed in claim 33 wherein the liquid comprises ammonium hydroxide.
- 5 41. A method for treating a workpiece as claimed in claim 33 wherein the step of spraying is further defined by spraying the liquid onto the workpiece from one or more fixed spraying positions.
- 10 42. A method for treating a workpiece as claimed in claim 41 and further comprising the step of rotating the workpiece as it is sprayed with the ozone and liquid mixture.
- 15 43. A method for treating a workpiece as claimed in claim reference 11 and further comprising the step of rotating the workpiece as it is sprayed with the ozone and heated liquid mixture.
- 20 44. An apparatus for supplying a mixture of a treatment liquid and ozone onto the surface of a workpiece, the apparatus comprising:
a liquid reservoir having a liquid chamber;
a pump having an input in fluid communication with the liquid chamber,
the pump further having an output;
one or more nozzles disposed to spray fluid therefrom onto the surface of the workpiece;

- a fluid path extending between the output of the pump and the one or more
nozzles, the fluid path carrying pressurized liquid provided at the
output of the pump;
an ozone supply system for injecting ozone into the fluid path; and
5 a heater disposed to heat the treatment liquid prior to supply of the
treatment liquid onto the surface of the workpiece.

45. An apparatus as claimed in claim 44 and further comprising a mixer disposed
in the fluid path.

10

46. An apparatus as claimed in claim 44 wherein the ozone supply system
comprises a contactor disposed to receive the ozone and the treatment liquid.

47. An apparatus as claimed in claim 45 wherein the mixer is a static mixer.

15

48. An apparatus as claimed in claim 45 wherein the mixer is an active mixer.

49. An apparatus as claimed in claim 44 wherein the pressurized liquid is
deionized water.

20

50. An apparatus as claimed in claim 44 wherein the pressurized liquid comprises
sulfuric acid.

51. An apparatus as claimed in claim 44 wherein the pressurized liquid comprises ammonium hydroxide.
52. An apparatus as claimed in claim 44 wherein the pressurized liquid comprises
5 an acid hydroxide.
53. An apparatus as claimed in claim 44 wherein the pressurized liquid comprises hydrochloric acid.
- 10 54. An apparatus as claimed in claim 44 wherein the liquid reservoir comprises an inlet for supplying ozone into a liquid chamber of the liquid reservoir, the liquid chamber being in fluid communication with the pump.
55. An apparatus as claimed in claim 54 wherein the inlet for supplying ozone is in
15 fluid communication with the ozone generator
56. An apparatus as claimed in claim 54 wherein the liquid reservoir further comprises an ozone dispersion apparatus disposed in the liquid chamber for dissolving ozone received at the inlet in liquid contained within the liquid
20 chamber.
57. An apparatus as claimed in claim 44 and further comprising a chamber for housing the workpiece as it is sprayed with treatment liquid that proceeds from the one or more nozzles.

58. An apparatus as claimed in claim 57 and further comprising a re-circulation fluid path extending between the chamber and the liquid reservoir.

5 59. An apparatus as claimed in claim 57 and further comprising a rotor assembly disposed in the chamber for rotating the workpiece.

60. An apparatus for providing a mixture of ozone and a liquid comprising water for treatment of the surface of a workpiece, the apparatus comprising:

10 a reservoir having a chamber holding the liquid comprising water;
a pump having an input connected to receive the liquid comprising water from the chamber of the reservoir, the pump further having an output for supplying pressurized liquid comprising water therefrom;
one or more nozzles disposed to spray fluid therefrom onto the surface of
15 the workpiece;
a fluid path extending between the output of the pump and the one or more nozzles, the fluid path carrying the pressurized liquid comprising water that is provided at the output of the pump; and
an ozone generator for generating a supply of ozone at an output thereof;
20 one or more supplying lines extending from the ozone generator to the fluid path for injecting ozone into the fluid path.

61. An apparatus as claimed in claim 60 and further comprising a mixer disposed in the fluid path.

62. An apparatus as claimed in claim 61 wherein the mixer is a static mixer.
63. An apparatus as claimed in claim 61 wherein the mixer is an active mixer.
- 5 64. An apparatus as claimed in claim 60 wherein the reservoir comprises an inlet
for supplying ozone into the chamber of the reservoir.
- 10 65. An apparatus as claimed in claim 64 wherein the inlet for supplying ozone is in
fluid communication with the ozone generator
- 15 66. An apparatus as claimed in claim 60 wherein the reservoir further comprises an
ozone dispersion apparatus disposed in the chamber for dissolving ozone
received at the inlet in the liquid comprising water contained within the
chamber.
- 20 67. An apparatus as claimed in claim 60 and further comprising a chamber for
housing the workpiece as it is sprayed with the liquid comprising water that
proceeds from the one or more nozzles.
68. An apparatus as claimed in claim 67 and further comprising a re-circulation
fluid path extending between the chamber housing a workpiece and the
chamber of the reservoir.

69. An apparatus as claimed in claim 67 and further comprising a rotor assembly disposed in the chamber for rotating the workpiece, the rotor assembly being operable up to at least 300 rpm.
- 5 70. An apparatus for supplying a mixture of a treatment liquid and ozone for treatment of a surface of a workpiece, the apparatus comprising:
- means for heating the workpiece;
 - means for spraying the treatment liquid onto the surface of the workpiece;
 - means for introducing ozone into an environment containing the workpiece;
 - 10 means for controlling the thickness of a boundary layer of the treatment liquid on the surface of the workpiece to thereby facilitate diffusion of the ozone to the surface of the workpiece.
71. An apparatus as claimed in claim 70 wherein the means for heating comprises a
- 15 heater disposed to heat the treatment liquid that is applied to the workpiece.
72. An apparatus as claimed in claim 70 wherein the means for controlling comprises a rotor disposed to rotate the workpiece.
- 20 73. An apparatus as claimed in claim 70 wherein the means for controlling comprises means for controlling the flow of treatment liquid applied to the surface of workpiece.

74. An apparatus as claimed in claim 70 wherein the means for controlling the flow of treatment liquid applied to the surface of the workpiece comprises a pump.
- 5 75. An apparatus as claimed in claim 70 wherein the means for controlling comprises one or more nozzles adapted to generate fine droplets of the treatment liquid.
- 10 76. An apparatus as claimed in claim 70 wherein the treatment liquid is deionized water.
77. An apparatus as claimed in claim 70 wherein the treatment liquid comprises an acid.
- 15 78. An apparatus as claimed in claim 70 wherein the treatment liquid comprises sulfuric acid.
79. An apparatus as claimed in claim 70 wherein the treatment liquid comprises hydrochloric acid.
- 20 80. An apparatus as claimed in claim 70 wherein the treatment liquid comprises ammonium hydroxide.

81. An apparatus as claimed in claim 76 wherein the means for heating comprises a steam boiler.
82. An apparatus as claimed in claim 70 wherein the treatment liquid is an aqueous solution comprising water.
83. An apparatus as claimed in claim 82 wherein the means for heating comprises a steam boiler.
84. An apparatus for supplying a mixture of a treatment liquid and ozone for treatment of a surface of a workpiece, the apparatus comprising:
 - a heater disposed to heat the workpiece;
 - a reservoir;
 - a treatment chamber housing the workpiece;
 - a liquid supply line providing fluid communication of the treatment liquid between the reservoir and the treatment chamber;
 - one or more nozzles accepting treatment liquid from the liquid supply line and spraying the treatment liquid onto the surface of the workpiece;
 - an ozone generator having an output line for providing ozone into an environment containing the workpiece;
 - means for controlling the thickness of a boundary layer of the heated treatment liquid on the surface of the workpiece to thereby facilitate diffusion of the ozone to the surface of the workpiece.

85. An apparatus as claimed in claim 84 wherein the heater indirectly heats the workpiece by heating the treatment liquid that is sprayed thereon.

5 86. An apparatus as claimed in claim 84 wherein the treatment liquid is deionized water.

87. An apparatus as claimed in claim 84 wherein the treatment liquid comprises an acid.

10 88. An apparatus as claimed in claim 84 wherein the treatment liquid comprises sulfuric acid.

89. An apparatus as claimed in claim 84 wherein the treatment liquid comprises hydrochloric acid.

15

90. An apparatus as claimed in claim 84 wherein the treatment liquid comprises ammonium hydroxide.

20 91. An apparatus as claimed in 84 wherein the treatment liquid is an aqueous solution comprising water.

92. An apparatus as claimed in claim 84 wherein the output line of the ozone generator is connected to the liquid supplying line.

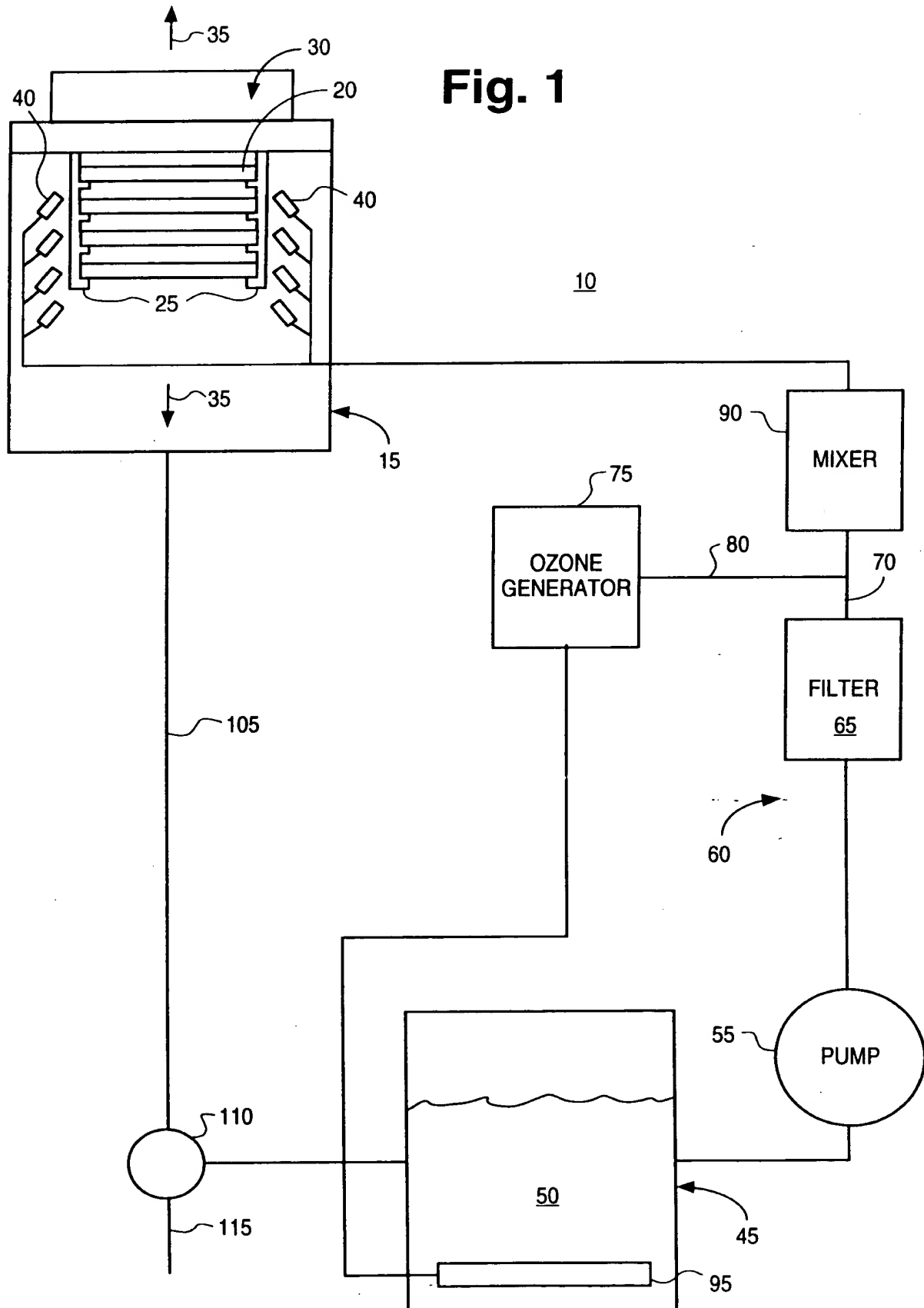
93. An apparatus as claimed in claim 84 wherein the output line of the ozone generator is connected to the treatment chamber.
94. An apparatus as claimed in claim 84 wherein the heater is connected in-line
5 with the liquid supplying line.
95. An apparatus as claimed in claim 84 wherein the heater is disposed in the reservoir.
- 10 96. An apparatus as claimed in claim 84 wherein the means for controlling comprises a rotor disposed to rotate the workpiece.
97. An apparatus as claimed in claim 84 wherein the means for controlling comprises means for controlling the flow of treatment liquid through the liquid
15 supply line.
98. An apparatus as claimed in claim 84 wherein the means for controlling the flow of treatment liquid through the liquid supply line comprises a pump.
- 20 99. An apparatus as claimed in claim 84 wherein the means for controlling comprises one or more nozzles adapted to generate fine droplets of the treatment liquid.

ABSTRACT

A novel chemistry and application technique used in a chemical process sequence to address the contamination concerns of the microelectronic fabrication industry is set forth. These concerns are namely: minimizing contamination for organics, particles, metal/ions, and silicon dioxide. Generally stated, cleaning of workpieces, such as semiconductor wafers, is accomplished by delivery of a chemical stream to the workpiece surface. Ozone is delivered either into the liquid process stream or into the process environment. The chemical stream, which may be in the form of a liquid or vapor, is applied to the wafer in a system which enables control of the boundary layer which forms on the workpiece surface. The chemical stream may include components such as ammonium hydroxide for simultaneous particle and organic removal, another chemical to raise the pH of the solution, or other chemical additives designed to accomplish one or more specific cleaning tasks. The particular application technique is configured to develop and control a boundary layer of the liquid on the wafer surface.

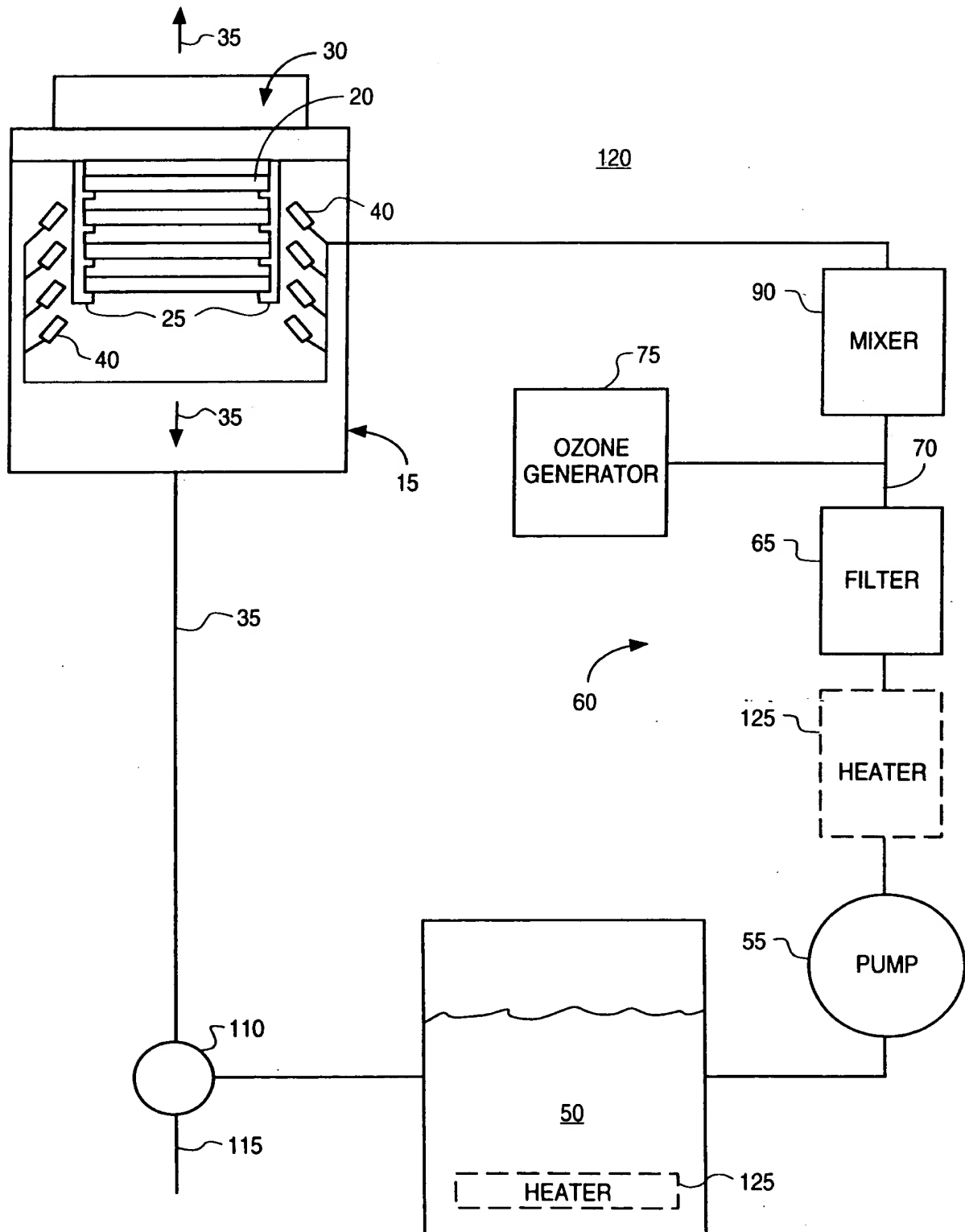
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Fig. 1



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Fig. 2



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Fig. 3

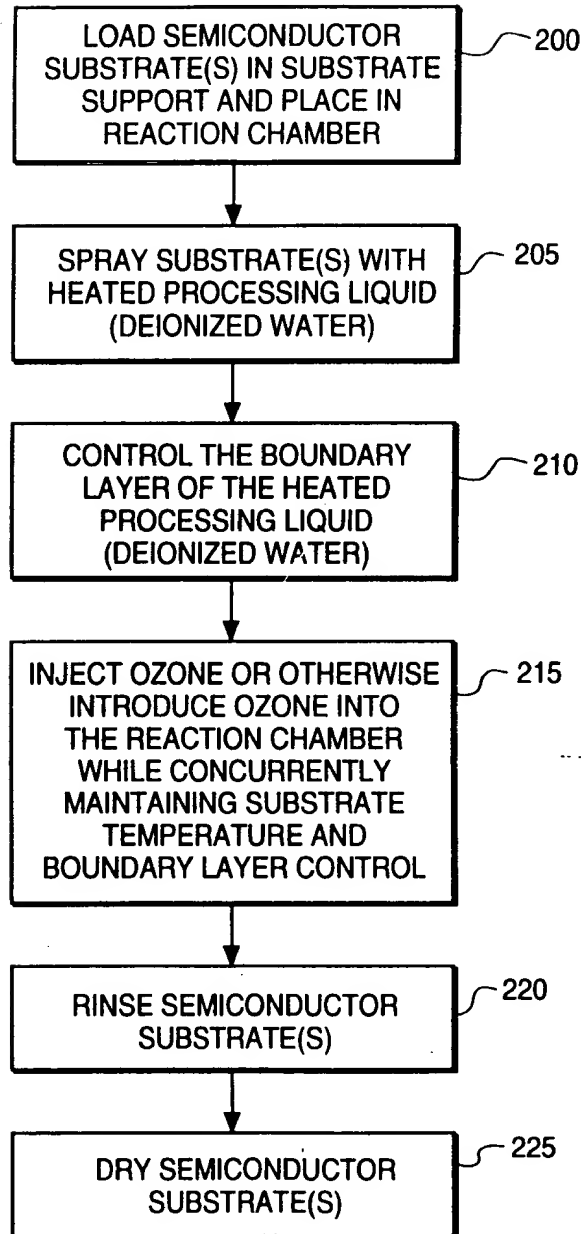
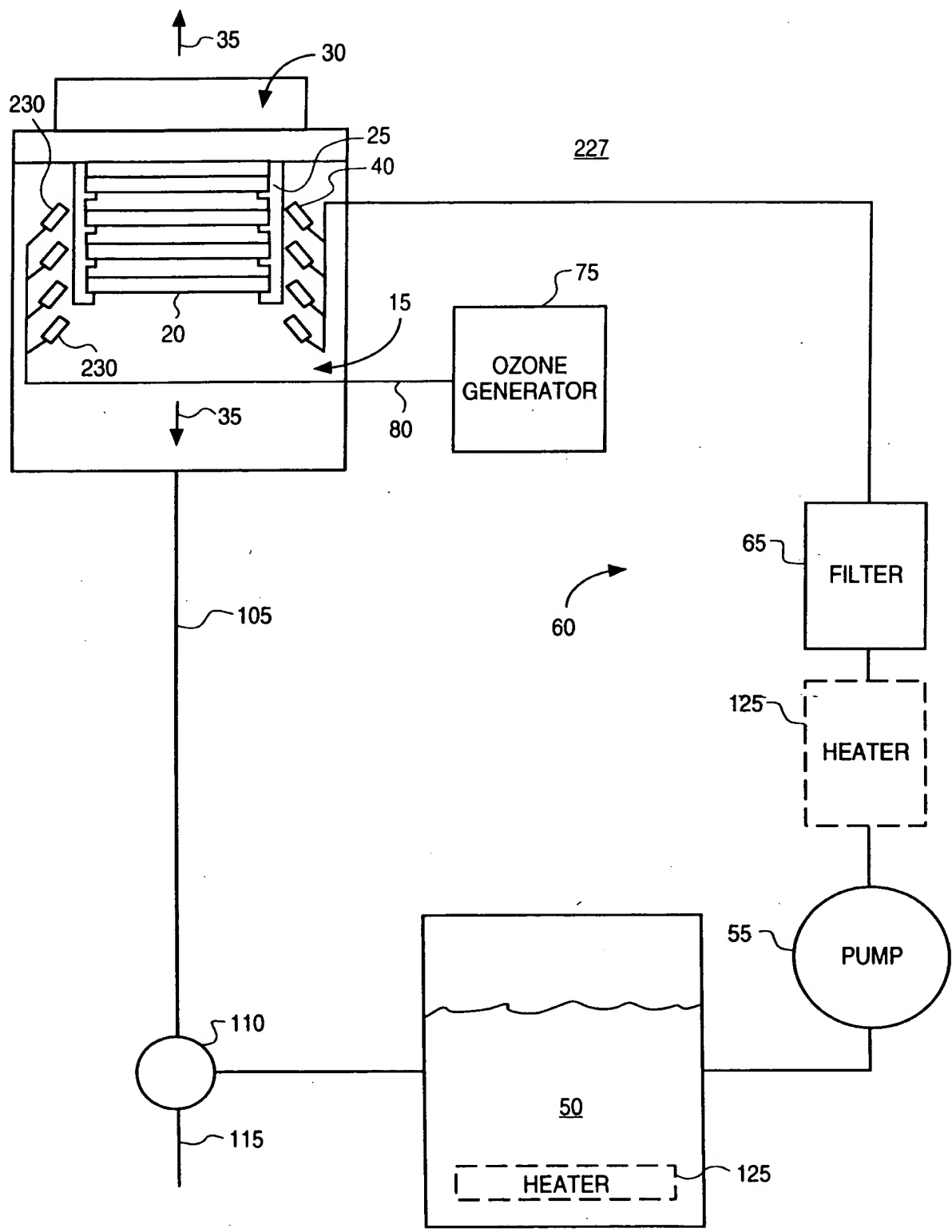
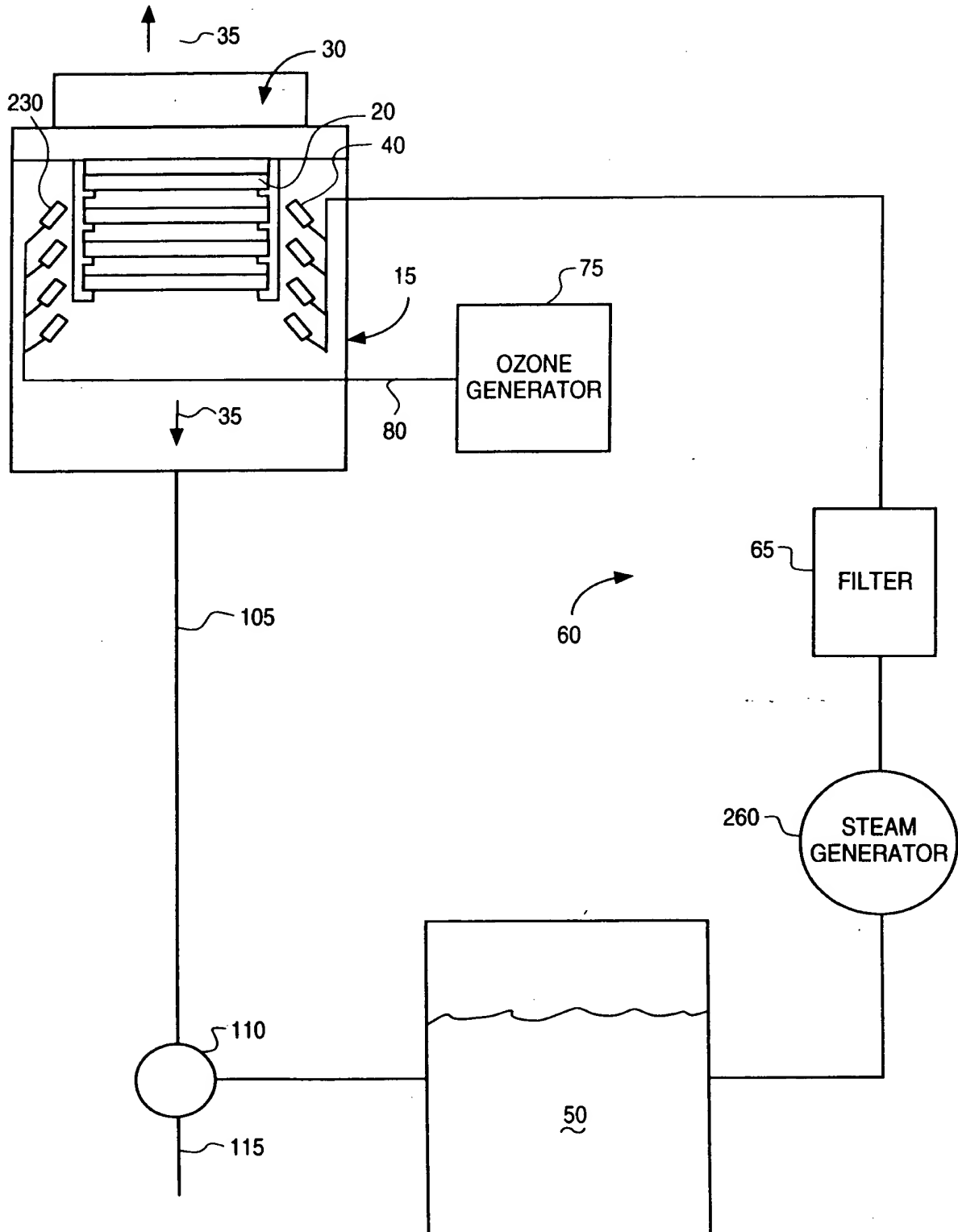


Fig. 4



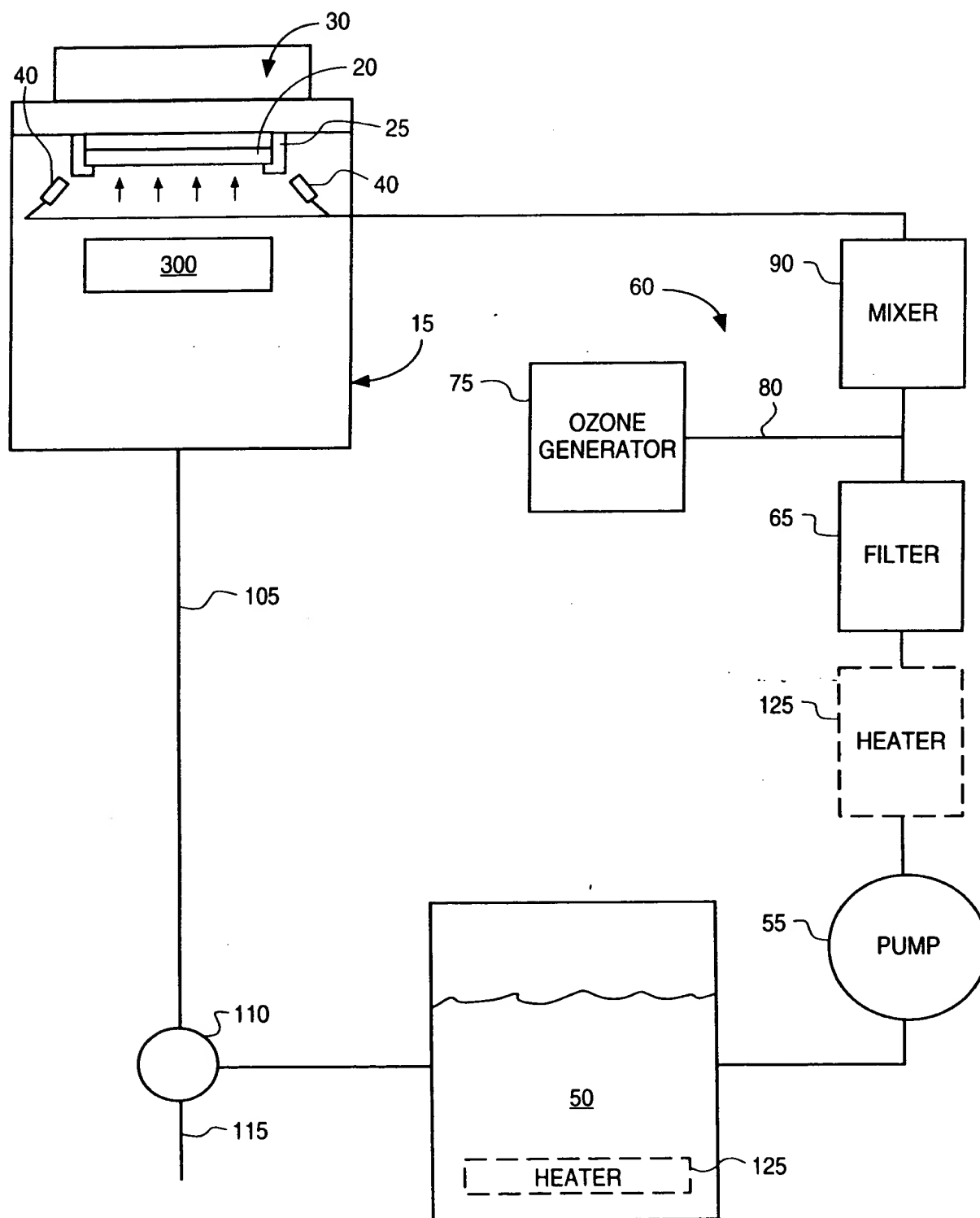
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Fig. 5



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Fig. 6



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Fig. 7

